```
(FILE 'HOME' ENTERED AT 20:42:37 ON 24 JUL 2007)
     FILE 'CA' ENTERED AT 20:42:54 ON 24 JUL 2007
    317116 S PHOSPHORYLAT? OR GLYCOSYLAT? OR DEACYLAT? OR ACYLAT?
\cdotL1
L2
    507353 S PHOSPHOKINASE OR KINASE OR TRANSFERASE OR PHOSPHORYLASE OR
            TRANSAMINASE OR AMINASE OR CARBOXYLASE OR CARBONYLASE OR
            PHOSPHATASE
    111473 S L1 AND L2
L3
L4
      3190 S L3 AND (FLUORESC? OR FLUORIMET? OR FLUOROMET?)
      3834 S L3 AND PEPTIDE AND SUBSTRATE
L5
L6
       255 S L4 AND L5
        61 S L6 AND PY<2000
L7
     29250 S L3 AND (PEPTIDE OR OLIGOPEPTIDE OR SUBSTRATE)
\Gamma8
L9
      1035 S L4 AND L8
L10
        87 S L9 AND QUENCH?
L11
       319 S L9 AND PY<2000
L12
       206 S L11 AND (HOMOGENEOUS OR ASSAY? OR DETERMIN? OR MICRODETERMIN? OR
            MEASUR? OR MONITOR? OR SENSE# OR SENSOR OR SENSING OR DETECT?) .
       287 S L7, L10, L12
L13
     FILE 'BIOSIS' ENTERED AT 21:10:37 ON 24 JUL 2007
L14
       213 S L13
     FILE 'MEDLINE' ENTERED AT 21:11:11 ON 24 JUL 2007
L15
       392 S L13
     FILE 'BIOSIS' ENTERED AT 21:13:12 ON 24 JUL 2007
       196 S L14 AND PY<2000
L16
     FILE 'MEDLINE' ENTERED AT 21:13:44 ON 24 JUL 2007
L17
       369 S L15 AND PY<2000
     FILE 'BIOSIS' ENTERED AT 21:17:15 ON 24 JUL 2007
L18
        85 S L7, L10
L19
        68 S L18 AND PY<2000
     FILE 'MEDLINE' ENTERED AT 21:18:33 ON 24 JUL 2007
       165 S L7, L10
L20
       142 S L20 AND PY<2000
L21
         8 S L17 NOT L21 AND (FLUORESC? OR FLUORIMET? OR FLUOROMET?) /TI
L22
     FILE 'CA, BIOSIS, MEDLINE' ENTERED AT 21:24:29 ON 24 JUL 2007
L23
       369 DUP REM L13 L19 L21 L22 (136 DUPLICATES REMOVED)
=> d bib, ab 123 1-369
L23
     ANSWER 51 OF 369 CA COPYRIGHT 2007 ACS on STN
     138:35759 CA
ΑN
TΙ
     Fluorescent protein sensors containing phosphorylation sites introduced
     by N-terminal mutagenesis
ΙN
     Cubitt, Andrew B.
     Aurora Biosciences Corporation, USA
PΑ
     U.S., 49 pp.
SO
     US 6495664
                          В1
                                 20021217
                                             US 1998-129192
                                                                    19980724
PΙ
PRAI US 1998-129192
                          A1
                                 19980724
AΒ
     The present invention includes a fluorescent compd. that can detect an
     activity, such as an enzymic activity, and exhibits quenching.
     fluorescent compd. is a fluorescent protein, such as an Aequorea-related
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green fluorescent protein.. The green fluorescent protein (GFP) of

Aequorea victoria is modified to include a substrate site for an enzymic activity such as a kinase activity, a phosphatase activity, a protease activity, and a glycosylase activity. Thus, relative fluorescence of phosphorylated vs. non-phosphorylated GFP is enhanced by modifying the N-terminal region (e.g., residues MSKGEELF to MGRRRASII) to contain a phosphorylation site responsive to protein kinase A, or other protein kinase enzymes,. Addnl. amino acid substitutions are engineered (S65A, K79R, E90N, N149K, V163A, I167T, and optionally A87T and E90A) to further improved fluorescence yield. The fluorescent compd. of the present invention can be used to detect such enzymic activities in samples, such as biol. samples, including cells. The present invention also includes nucleic acids that encode the fluorescent compds. of the present inventions, and cells that include such nucleic acids or fluorescent compds.

- L23 ANSWER 53 OF 369 CA COPYRIGHT 2007 ACS on STN
- ΑN 134:128210 CA
- TIHomogeneous fluorescence method for assaying structural modifications of biomolecules using double-labeled substrates

WO 2000-US40495

20000727

20010201

- IN Blumenthal, Donald K., II
- University of Utah Research Foundation, USA PA
- SO PCT Int. Appl., 35 pp.
- PΙ WO 2001007638 A2
- PRAI US 1999-145755P Р 19990727
- Double-labeled protein biomol. substrates and methods for the AB homogeneous assay of processes by which biomols. are covalently modified are described. The methods of the present invention utilize biomol. substrates labeled at two positions with two fluorescent dyes or with a **fluorescent** dye and a nonfluorescent dye. The two labeling dyes of the unmodified biomol. substrates stack, thereby quenching the substrate's fluorescence. Upon covalent modification of the double-labeled substrate, however, the intramolecularly stacked dyes dissoc. and the fluorescence of the phosphorylated substrate changes markedly. utilizing the double-labeled substrates of the present invention do not require phys. sepn. of modified and unmodified substrate mols., nor do they require other special reagents or radioactive materials. Methods for prepg. and characterizing the substrates used in the assay procedure are described, as are methods utilizing the substrates of the present invention for high-throughput screening, for monitoring intracellular processes of covalent biomol. modification in living cells, for diagnostic and therapeutic applications for diseases involving dysfunctional processes of covalent biomol. modification, and for discovering novel enzymic substrates. A synthetic KID peptide was prepd. and double-labeled with tetramethylrhodamine-5-maleimide and 5carboxyfluorescein, succinimidyl ester or 5-carboxytetramethylrhodamine, succinimidyl ester. These substrates can be used to assay for protein kinase A as the phosphorylated substrates have detectable changes in the absorbance and fluorescence characteristics of the dyes included in the substrates.

- TI Preparation of **peptides** having **fluorescence** and **fluorescence-quenching** groups as **substrates** for **determination** of protein **phosphatase**
- IN Nishikata, Makoto
- PA Japan
- SO Jpn. Kokai Tokkyo Koho, 11 pp.
- PI JP 11012297 A 19990119 JP 1998-126684 19980421 US 5917012 A 19990629 US 1998-70756 19980430
- PRAI JP 1997-126463 A 19970430
- AB (un)protected **peptides** represented by formula A1-X-A3 [X = PO3H2introduced amino acid residue; A1, A2 = (un)protected amino acid or peptide residue linked to X; either of A1 or A2 possesses a fluorescence-quenching group at the terminus or side-chain and the other possesses a fluorescent group possessing fluorescence property by its self at the terminus or side-chain with its fluorescence property being quenched by a fluorescence-quenching group in the mol. | are prepd. method for detn. of protein phosphatase comprises reaction of the above peptide deriv. with a protein phosphatase, reaction of the reaction product with protease, and measurement of the change in fluorescence A reagent for detn. of protease phosphatase activity contains the above peptide deriv. and protease which selectively cleaves the peptide bond between the fluorescence-quenching group-contg. amino acid residue and the **fluorescence** group-linked amino acid residue after dephosphorylation from PO3H2-introduced amino acid residue by protease phosphatase. The advantages of these peptide substrates are (1) protein phosphatase activity is measured in high sensitivity, since the assay measures fluorescence intensity, (2) they can be stably stored for a long period of time, since they do not contain radio active phosphorus compds., (3) the assay can be applied to a sample contg. phosphate ions, since it does not measure released phosphate ions, (4) a crude tissue ext. liq. can be also used as a sample, since the excitation wavelength is set to the wavelength different from the absorption frequency of proteins by appropriately choosing the **fluorescence** group, and (5) dephosphorylation reaction of the substrates can be traced in real time as long as protein phosphatase to detd. is resistant to protease having the above property. Thus, Mca-Gly-Glu-Gly-Thr-Tyr(PO3H2)-Gly-Lys(DNP)-Arg-NH2 (DNP = 2,4-dinitrophenyl, Mca = 7-methoxycumarin-4-ylacetyl),which was prepd. by the solid phase method using  $N\alpha$ -Fmoc-protected amino acids and a MBHA resin, enabled detn. of protein tyrosine phosphatase in real time with good accuracy.

## L23 ANSWER 94 OF 369 MEDLINE on STN

- AN 1999441240 MEDLINE
- DN PubMed ID: 10510304
- TI A phosphotyrosine-containing quenched fluorogenic peptide as a novel substrate for protein tyrosine phosphatases.
- AU Nishikata M; Suzuki K; Yoshimura Y; Deyama Y; Matsumoto A
- CS Department of Dental Pharmacology, Hokkaido University School of Dentistry, Sapporo 060-0813, Japan.. mnishika@den.hokudai.ac.jp
- SO The Biochemical journal, (1999 Oct 15) Vol. 343 Pt 2, pp. 385-91.
- AB Mca-Gly-Asp-Ala-Glu-Tyr(PO(3)H(2))-Ala- Ala-Lys(DNP)-Arg-NH(2), where Mca is (7-methoxycoumarin-4-yl)acetyl and DNP is 2,4-dinitrophenyl, was synthesized as a fluorogenic **substrate** for protein tyrosine **phosphatases**

(PTPs). In the peptide, the fluorescent Mca group is quenched efficiently by the DNP group. Although the fluorescence intensity of the substrate was practically unchanged upon PTP-catalysed dephosphorylation, it increased approx. 120-fold upon subsequent treatment with chymotrypsin. Analysis by HPLC showed that chymotrypsin cleaved only the dephosphorylated substrate at the Tyr-Ala bond. with the aid of chymotrypsin, dephosphorylation of the substrate can be measured fluorometrically. A strictly linear correlation was observed between PTP concentration and dephosphorylation rate. The fluorogenic substrate was dephosphorylated by some PTPs much more rapidly than the corresponding (32) P-labelled substrate used for comparison, whereas alkaline phosphatase dephosphorylated the two substrates at similar The fluorogenic substrate is therefore more specific for PTPs than the radiolabelled **substrate**. The assay with the fluorogenic substrate could be applied to the estimation of kinetc parameters and measurement of PTP activity in crude-enzyme preparations. detection limit of our assay (1 microM substrate in 200 microliter of reaction mixture) was estimated to be 0.2-0.4 pmol, whereas it was estimated to be about 1 pmol in the assay that used (32)P-labelled peptide (specific radioactivity of approx. 1000 c.p.m. /pmol). Our assay is simple, specific, highly sensitive and non-radioisotopic, and hence would contribute greatly to the development of PTP biology.

- L23 ANSWER 109 OF 369 CA COPYRIGHT 2007 ACS on STN
- AN 128:151089 CA
- TI Assays for protein kinases using fluorescent protein substrates
- IN Tsien, Roger Y.; Cubitt, Andrew B.
- PA Regents of the University of California, USA; Tsien, Roger Y.; Cubitt, Andrew B.
- SO PCT Int. Appl., 69 pp.
- PI WO 9802571 A1 19980122 WO 1997-US12410 19970716 US 5912137 A 19990615 US 1996-679865 19960716
- PRAI US 1996-679865 A1 19960716
- This invention provides assays for protein kinase activity using fluorescent proteins engineered to include sequences that can be phosphorylated by protein kinases. The proteins exhibit different fluorescent properties in the non-phosphorylated and phosphorylated states. Aequorea victoria green fluorescent proteins contg. substitution mutations were prepd. with recombinant Escherichia coli. Some displayed increased fluorescence upon phosphorylation, other decreased fluorescence.
- L23 ANSWER 269 OF 369 CA COPYRIGHT 2007 ACS on STN
- AN 117:247747 CA
- TI Measurement of enzyme activity with labelled peptide substrates
- IN Ikenaka, Tokuji; Mega, Tomohiro; Hamazume, Yasuki
- PA Wako Pure Chemical Industries, Ltd., Japan
- SO U.S., 12 pp. Cont.-in-part of U.S. Ser. No. 32,253, abandoned.
- PI US 5120644 A 19920609 US 1988-256078 19881006
- PRAI JP 1986-76349 A 19860401
- AB Enzymes that modify proteins are assayed using peptide substrates with a fluorescent label. Enzymes that are assayed include endopeptidases,

proteinases, transferases, kinases, and phosphatases. The synthesis of the renin substrate 2-pyridyl-Pro-Phe-His-Leu-Val-Tyr- $\beta$ -Ala from 2-pyridyl glycine and BOC amino acids is described. This was used as an assay substrate for renin with the **fluorescent** cleavage product clearly separable from the substrate by HPLC.

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=> d his

(FILE 'HOME' ENTERED AT 19:04:38 ON 24 JUL 2007) FILE 'CA' ENTERED AT 19:04:50 ON 24 JUL 2007

- L1 1 S DICKENS ?/AU AND BIOCHEM?/SO AND BIOPHYS?/SO AND 1991/PY
- L2 0 S ROSSOMONDO ?/AU AND 1992/PY
- L3 1 S ROSSOMONDO ?/AU
- L4 10 S MITOGEN AND KINASE AND 1992
- L5 181 S MITOGEN AND KINASE AND 1992/PY
- L6 16 S L5 AND SCI?/SO
- L7 9 S L6 AND NAT?/SO
- L8 13 S PEARSON ?/AU AND BIOL?/SO AND 1985/PY
- L9 2 S L8 AND CHEM?/SO
- L10 2 S L7 AND ROSSOMANDO ?/AU
- L11 5 S L1, L9-10
- => d bib, ab, it 1-5
- L11 ANSWER 2 OF 5 CA COPYRIGHT 2007 ACS on STN
- AN 117:85711 CA
- TI The phorbol ester-dependent activator of the **mitogen**-activated protein **kinase** p42mapk is a **kinase** with specificity for the threonine and tyrosine regulatory sites
- AU Rossomando, Anthony; Wu, Jie; Weber, Michael J.; Sturgill, Thomas W.
- CS Dep. Intern. Med., Univ. Virginia, Charlottesville, VA, 22908, USA
- SO Proceedings of the National Academy of Sciences of the United States of America (1992), 89(12), 5221-5
- AB Mitogen-activated protein kinases (MAP kinases) are activated by dual tyrosine and threonine phosphorylations in response to various stimuli, including phorbol esters. To define the mechanism of activation, recombinant wild-type 42-kDa MAP kinase (p42mapk) and a kinase-defective mutant of p42mapk (K52R) were used to assay both activator activity for p42mapk and kinase activity toward K52R in stimulated EL4.IL2 mouse thymoma cells. Phorbol 12,13-dibutyrate (10 min, 650 nM) stimulated a single peak of MAP kinase activator that was coeluted from Mono Q at pH 7.5 and 8.9 with K52R kinase activity. Both activities were inactivated by the serine/threonine-specific phosphoprotein phosphatase 2A but not by the tyrosine-specific phosphoprotein phosphatase CD45. Phosphorylation of K52R occurred specifically on Thr-183 and Tyr-185, as detd. by tryptic phosphopeptide mapping in comparison with synthetic marker phosphopeptides. These findings indicated that phorbol esterstimulated MAP kinase kinase can activate p42mapk by threonine and tyrosine phosphorylations, and that p42mapk thus does not require an autophosphorylation reaction.

- L11 ANSWER 3 OF 5 CA COPYRIGHT 2007 ACS on STN
- ΑN 114:224317 CA
- ΤI Phosphorylation of tyrosines 1158, 1162 and 1163 on a synthetic dodecapeptide by the insulin receptor protein-tyrosine kinase
- ΑU Dickens, Martin; Tavare, Jeremy M.; Clack, Beatrice; Ellis, Leland; Denton, Richard M.
- CS Sch. Med. Sci., Univ. Bristol, Bristol, BS8 1TD, UK
- Biochemical and Biophysical Research Communications (1991), 174(2), 772-SO
- AΒ To investigate the mechanism of tyrosine phosphorylation by insulin receptor kinase, a synthetic dodecapeptide substrate (RRDIYETDYYRK; amino acids 1155-1165) contq. the 3 major insulin receptor autophosphorylation sites was utilized. All 3 tyrosines on this peptide were rapidly phosphorylated, and phosphorylation was probably initiated at Tyr-9. This peptide thus serves as a useful tool to study the mechanism of transphosphorylation by the insulin receptor. proteolytic activity was detected in purified receptor prepns. that removed basic residues from the peptide and prevented its binding to phosphocellulose paper. Such activity could pose a serious problem when using peptide substrates to assay for protein kinases in other acellular systems.
- ANSWER 4 OF 5 CA COPYRIGHT 2007 ACS on STN L11
- ΑN 103:209706 CA
- ΤI Substrate specificity of a multifunctional calmodulin-dependent protein kinase
- ΑU Pearson, Richard B.; Woodgett, James R.; Cohen, Philip; Kemp, Bruce E.
- CS Repatriation Gen. Hosp., Univ. Melbourne, Heidelberg, 3081, Australia
- SO
- Journal of Biological Chemistry (1985), 260(27), 14471-6 AB The substrate specificity of the multifunctional calmodulin-dependent protein kinase from skeletal muscle was studied by using a series of synthetic peptide analogs. The enzyme phosphorylated a synthetic peptide corresponding to the N-terminal 10 residues of glycogen synthase (Pro-Leu-Ser-Arg-Thr-Leu-Ser-Val-Ser-Ser-NH2), stoichiometrically at serine (Ser)-7, the same residue phosphorylated in the parent protein. The synthetic peptide was phosphorylated with a Vmax of 12.5 umol/min/mg and an apparent Km of 7.5  $\mu$ M compared to values of 1.2  $\mu$ mol/min/mg and 3.1 µM, resp., for glycogen synthase. Similarly, a synthetic peptide corresponding to the N-terminal 23 residues of smooth muscle myosin light chain was readily phosphorylated on Ser-19 with a Km of 4 uM and a Vmax of 5.4  $\mu$ mol/min/mg. The importance of the arginine found 3 residues N-terminal to the phosphorylated serine in each of these peptides was evident from expts. in which this arginine was substituted by either leucine or alanine, as well as from expts. in which its position in the myosin light chain sequence was varied. arginine-16 at residues 14 or 17 abolished phosphorylation, whereas location at residue 15 not only decreased Vmax 14-fold but switched the major site of phosphorylation from Ser-19 to threonine-18. The sequence 'Arg-X-Y-Ser(Thr) (where X and Y are any amino acids) apparently represents the min. specificity determinant for the multifunctional calmodulin-dependent protein kinases. Studies with various synthetic peptide substrates and their analogs revealed that the specificity determinants of the multifunctional calmodulin-dependent protein kinase

were distinct from several other arginine-requiring protein kinases.

- L11 ANSWER 5 OF 5 CA COPYRIGHT 2007 ACS on STN
- AN 103:33969 CA
- TI Spatial requirements for location of basic residues in peptide substrates for smooth muscle myosin light chain kinase
- AU Kemp, Bruce E.; Pearson, Richard B.
- CS Dep. Med., Univ. Melbourne, Heidelberg, 3081, Australia
- SO Journal of Biological Chemistry (1985), 260(6), 3355-9
- The requirement of basic residues as substrate specificity determinants AΒ for chicken gizzard myosin light-chain kinase (I) was studied by using synthetic peptide analogs of the local phosphorylation site sequence in the myosin light chains, Lys-Lys-Arg13-Pro-Gln-Arg16-Ala-Thr-Ser19-Asn-Val-Phe-Ala. The basic residue arginine (Arg)-16, exerted a strong influence on the kinetics of phosphorylation similar to that reported previously for the 3-adjacent residues, lysine (Lys)-11, Lys-12, and The location of Arg-16 in relation to serine (Ser)-19 as well as the distance between Arg-13 and Arg-16 had a profound effect on both the kinetics and the site specificity of phosphorylation. Placement of Arg-16 at position 15 resulted in a complete switch in phosphorylation site specificity from Ser-19 to threonine (Thr)-18. Increasing the no. of alanine residues between Arg-13 and Arg-16 in the model peptide, Lys-Lys-Arg-(Ala)n-Arg-Ala-Thr-Ser-Asn-Val-Phe-Ala, also influenced the kinetics and site specificity of peptide phosphorylation. With 2 or 3 alanines (n = 2 or 3), the apparent Km was 7.5 and 10  $\mu$ M, resp., and 97% of the phosphate was esterified to Ser-19. Increasing or decreasing the no. of alanines (n = 0 to n = 4) was accompanied by an increase in the apparent Km and phosphorylation of both Thr-18 and Ser-19. results supported the concept that both the presence and location of basic residues play an essential role in the substrate specificity of the smooth muscle I.

=> log y STN INTERNATIONAL LOGOFF AT 19:24:50 ON 24 JUL 2007